

DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service Food and Drug Administration SOUTHWEST REGION

Office of the Regional Food and Drug Director 7920 Elmbrook Drive, Suite 102 Dallas, TX 75247-4982 TELEPHONE: 214-655-8100 FACSIMILE: 214-655-8130

June 15, 2001

WARNING LETTER

CERTIFIED MAIL RETURN RECEIPT REQUESTED

01-SWR-WL-54/F

Wayne Masiker NCOIC Radiology Department 67th Combat Support Hospital Unit 26610 APO, AE 09244

RE: Inspection ID - 2049580007

Dear NCOIC Wayne Masiker,

On May 23, 2001, a representative of the Food and Drug Administration (FDA) inspected your facility. This inspection revealed a serious regulatory problem involving the mammography at your facility.

The Mammography Quality Standards Act of 1992 requires your facility to meet specific standards. These requirements help protect the health of women by assuring that a facility can perform quality mammography. The inspection revealed the following level 1 finding at your facility:

Level 1: Mammograms were processed in processor 1, Kodak, when it was out of limits on at least 5 days.

The specific problem noted above appeared on your MQSA Facility Inspection Report, which was issued to your facility at the close of the inspection.

Level 1 findings may be symptomatic of serious underlying problems that could compromise the quality of mammography at your facility. They represent a serious violation of the law which may result in FDA taking regulatory action without further notice to you. These actions include, but are not limited to:

- Placing your facility under a Directed Plan of Correction.
- Charging your facility for the cost of on-site monitoring.
- Assessing civil money penalties up to \$10,000 for each failure to substantially comply with, or each day of failure to substantially comply with, the Standards.

- Suspension or revocation of your facility's FDA certificate, or obtaining a court injunction against further mammography.

In addition, your response should address the level 2 finding that was listed on the inspection report provided to you at the close of the inspection. The inspection revealed the following level 2 finding:

Level 2: Corrective actions for processor QC failures were not documented at least once for processor 1, Kodak.

It is necessary for you to act on this matter immediately. You are required to respond to this office in writing within fifteen (15) working days from receipt of this letter. Please address the following:

- The specific steps you have taken to correct all of the violations noted in this letter.
- Each step your facility is taking to prevent the recurrence of similar violations.
- Equipment settings (including technique factors), raw test data, and calculated final results, where appropriate.
- Sample records that demonstrate proper record keeping procedures, if the findings relate to quality control or other records (Note: Patient names or identification should be deleted from any copies submitted).

Please submit your response to: Deborah M. McGee, Radiation Specialist Food and Drug Administration 7920 Elmbrook Drive, Suite 102 Dallas, TX 75247-4982

This letter pertains only to findings of your inspection and does not necessarily address other obligations you have under the law. You may obtain general information about all of FDA's requirements for mammography facilities by contacting the Mammography Quality Assurance Program, Food and Drug Administration, P.O. Box 6057, Columbia, MD 21045-6057 (1-800-838-7715) or through the Internet at http://www.fda.gov.

If you have more specific questions about mammography facility requirements, or about the content of this letter, please feel free to contact Deborah M. McGee at (214) 655-8100 ext. 138.

Sincerely yours,

Gary L. Pierce

Regional Food and Drug Director